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(54) THE: GENE ENCODING ALKALINE LIQUEFYING ALPHA-AMYLASE

(57) Abstract

The present invention provides a DNA fragment encoding alkaline liquefying α -amytase, recombinant DNA containing the DNA fragment, a transformed microorganism harboring the recombinant DNA, as well as a method for producing alkaline liquefying α -amytase using the transformant. The method of the present invention enables mass production of alkaline liquefying α -amylase useful as a detergent companient.

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Description

Gene Encoding Alkaline Liquefying Alpha-Amylase

Technical Field:

The present invention relates to the gene encoding alkaline liquefying α -amylase and fragments thereof, and to recombinant DNA and a transformant bearing the gene or fragments of the gene.

Background Art:

Alpha-amylase has long been used in a variety of fields. For example, it has been used for the saccharification of grains and potatoes in the fermentation industry, as starch paste removers in the textile industry, as digestives in the pharmaceutical industry, and for the manufacture of thick malt syrups in the food industry. Alpha-amylase is an enzyme which acts on a starch-related polysaccharides such as amylose and amylopectin, hydrolyzing solely the α -1,4-glucoside bond of the polysaccharide molecule. Since 1833, when Payen and Persoz first discovered the enzyme, crystalline samples or electrophoretically homogeneous samples of α -amylase have been obtained from a number of different sources including bacteria, fungi, plant seeds, and animal digestive glands.

The present inventors have recently discovered that the efficacy of dish-washing detergents and laundry detergents for clothes can be greatly improved, particularly on starch

dirts, when a-amylase and a debranching enzyme are both incorporated into these detergents (Japanese Fatent Application Laid-open (kokai) No. 2-132192). However, most of the a-amylases previously found in the natural world exhibit maximal and stable enzymatic activities in the neutral to acidic pH ranges, but scarcely work in an alkaline solution of pH 9-10. There exist only a small number of amylase enzymes that are known to exhibit maximal activities in the alkaline pH range (so-called alkaline lphaamylases and alkali-resistant α -amylases). These alkaline α amylases and alkali-resistant q-amylase include, an enzyme produced by Bacillus sp. A-40-2 [Horikoshi, K. et al., Agric. Biol. Chem., 35, 1783 (1971)], an enzyme produced by Bacillus sp. NRRL B-3881 (Boyer, E., J. Bacteriol., 110, 992 (1972)], an enzyme produced by Streptomyces sp. KSM-9 (Japanese Patent Application Laid-Open (kokai) No. 61-209528, an enzyme produced by Bacillus sp. H-167 (Japanese Patent Application Laid-Open (kokai) No. 62-208278, an enzyme produced by Bacillus alkalothermophilus A3-8 (Japanese Patent Application Laid-Open (kokai) No. 2-49584, and an enzyme produced by Natronococcus sp. Ah-36 (Japanese Patent Application Laid-Open (kokai) No. 4-211369.

As used herein, the term "alkaline α -amylase" refers to α -amylases whose optimum pHs fall within the alkaline pH range, whereas the term "alkali-resistant α -amylase" refers to α -amylases which have optimum pHs within the neutral to acidic range but whose activities in the alkaline range are comparable with those obtained at an optimum pH, and in

addition, which retain their stabilities in the alkaline range. By the term "neutral range" is meant the range of pH not less than 6 and less than 8, and the term "alkaline" denotes a pH which is higher than the "neutral range".

Most of these alkaline α -amylases and alkali-resistant amylases are so-called saccharifying a-amylases which decompose starch or starch-related polysaccharides to glucose, maltose, or maltotriose. As such, these enzymes cause problems if they are used as enzymes for detergents, though they are advantageously used in the manufacture of sugar. Thus, there remains a need for so-called alkaline liquefying a-amylases which exhibit resistance against surfactants used in detergents, and which decompose starch or starchrelated polysaccharides in a highly random manner. The present inventors continued an extensive search for microorganisms producing an alkaline liquefying a-amylase suitable as a detergent component, and they discovered that an alkalophilic Bacillus sp. KSM-AP1378 strain, having its optimum pH for growth in the alkaline range, produces an enzyme exhibiting the activity of an alkaline liquefying a-amylase. They elucidated that this enzyme is useful as an additive in detergent compositions for washing dishes and kitchen utensils and for detergent compositions for clothes (WO94/26881).

Amounts of the enzyme produced may be effectively increased by improving a method for culturing an alkaline liquefying a-amylase-producing microorganism, Bacillus sp. KSM-AP1378, or by exploiting mutation. However, in order to

produce the enzyme advantageously on an industrial scale, another approach must be taken.

Amounts of an enzyme produced can be enhanced using a genetic engineering approach, and in addition, the catalytic properties of the enzyme can be improved, using a protein engineering approach, by altering the gene encoding the enzyme. However, the gene encoding an alkaline liquefying α -amylase has not yet been obtained.

Accordingly, an object of the present invention is to provide the gene encoding alkaline liquefying α -amylase and fragments thereof, a transformant harboring recombinant DNA comprising the gene, and a method for producing an alkaline liquefying α -amylase using the transformant.

The DNA encoding the alkaline liquefying α -amylase gene may be further used to produce probes to be used in the isolation of additional, homologous alkaline liquefying α -amylase genes from other microorganisms. Thus, an additional object of the present invention is to provide a means of screening for and isolating additional alkaline liquefying α -amylase enzymes.

Disclosure of the Invention

The present inventors attempted to isolate, from the chromosomal DNA of an alkalophilic Bacillus strain, a DNA fragment containing the gene encoding an alkaline liquefying a-amylase, and as a result, they were successful in isolating an approximately 1.8 kb DNA fragment encoding an alkaline liquefying a-amylase. When they transformed a host microorganism using this DNA fragment ligated to a

suitable vector, it was confirmed that the resultant recombinant microorganism produced an alkaline liquefying a-amylase. Moreover, it was found that the amino acid sequence of the alkaline liquefying a-amylase to be encoded is different from that of previously known amylases. The present invention was accomplished based on this finding.

Accordingly, the present invention provides a DNA fragment encoding an alkaline liquefying $\alpha\text{-amylase}$.

The present invention also provides a recombinant DNA comprising the above-described DNA fragment encoding an alkaline liquefying α -amylase.

The present invention also provides a transformed microorganism harboring the above-described recombinant DNA comprising a DNA fragment encoding an alkaline liquefying a-amylase.

The present invention further provides a method for producing an alkaline liquefying a-amylase, by culturing the above-described transformed microorganism and collecting the enzyme.

Brief Description of the Drawings

Fig. 1 shows a restriction enzyme map of a fragment of the gene encoding an alkaline liquefying amylase;

Fig. 2 is a chart depicting construction of pAML100 using a fragment of the gene encoding an alkaline liquefying amylase:

Fig. 3 shows nucleotide sequences of primers used.

Fig. 4 is a pH profile of an alkaline liquefying $\alpha-$ amylase produced by Bacillus sp. KSM-AP1378.

Best Mode for Carrying Out the Invention

In the present invention, a useful microorganism which serves as an alkaline liquefying \$\alpha\$-amylase gene donor may be, for example, Bacillus sp. KSM-AP1378 (FERM BP-3048, deposited July 24, 1989 in Fermentation Research Institute, Agency of Industrial Science and Technology of 1-3, Higashi 1-chome, Tsukuba-shi, Ibaraki, 305 Japan), which is an alkalophilic Bacillus strain. This strain was isolated from the soil in the vicinity of the city of Tochigi in Tochigi Prefecture, Japan by the present inventors and identified as a strain which produces significant amounts of alkaline liquefying \$\alpha\$-amylase. This strain was deposited at the Fermentation Research Institute, Agency of Industrial Science and Technology (1-3, Higashi 1-chome, Tsukuba-shi, Ibaraki-ken, 305, Japan) under FERM BP-3048 on August 8, 1990 (originally deposited as P-10885 on July 24, 1989).

In order to obtain chromosomal DNA from a donor microorganism, the method proposed by Marmur, J. (<u>J. Mol. Biol.</u>, 3, 208 (1961)) or the method proposed by Saito, H. and Miura, K. (<u>Biochem. Biophys. Acta</u>, 72, 619 (1963)) may be used. Other similar methods may also be used.

DNA fragments comprising the alkaline liquefying a-amylase gene are prepared by cleaving the thusobtained chromosomal DNA using restriction enzymes. Restriction enzymes which may be used are not particularly

limited so long as they do not fragment the gene. alkaline liquefying a-amylase gene may also be obtained by PCR (Mullis, K.B. and Falcona, F.A., Methods Enzymol., 155, 335 (1987): Saiki, R. K. et al., Science, 239, 487 (1988). For example, the gene may be obtained through the synthesis of primers having sequences corresponding to those on the upstream side of the 5'terminus and on the downstream side of the 3'-terminus of the essential region based on the nucleotide sequence described in Sequence No. 2, and subsequently conducting PCR using, the chromosomal DNA of an alkaline liquefying a-amylase-producing microorganism as a template. Alternatively, an intact gene may be obtained by first obtaining a fragment of the alkaline liquefying a-amylase gene from an alkaline liquefying a-amylaseproducing microorganism using any procedure, followed by PCR which amplifies the upstream and downstream sides of the fragmentary gene.

The thus-prepared genetic fragment is then subjected to cloning. Host/vector systems which may be used are not particularly limited, so far as that host bacterial strains express the alkaline liquefying a-amylase gene of the present invention, that the recombinant DNA molecules can be replicated in the host bacteria, and that the integrated gene can be stably harbored. For example, members of the EK system in which the host is E. coli K-12, and members of the BS system in which the host is Bacillus subtilis Marburg, may be used. Use of the EK system, which

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encompasses many kinds of vectors and which is extensively studied genetically, provides good results and thus is preferred. Specific examples of host bacteria include HB101, C600, and JM109 of the EK system, and BD170, MI112, and ISW1214 of the BS system. Specific examples of vectors include pBR322 and pUC18 for the EK system, and pUB110 and pHY300PLK for the BS system.

A recombinant plasmid DNA molecule is created by cleaving a vector with a restriction enzyme followed by ligation with the above-mentioned chromosomal or PCR-amplified DNA fragment. The ligation may be achieved, for example, through the use of a DNA ligase.

Methods for transforming host bacterial strains using a recombinant DNA molecule are not particularly limited. For example, a calcium chloride method (Mandel, M. and Higa, A., J. Mol. Biol., 53, 159 (1970)) may be used in the case of hosts of the EK system, and a protoplast method (Chang, C. and Cohen, S.N., Mol. Gen. Genet., 168, 111 (1978)) may be used in the case of hosts of the BS system. Selection of recombinant microorganisms are performed as follows. First, microorganisms which have been transformed with DNA which contains a vector-derived DNA fragment are selected, using as an index a character which is not inactivated by insertion of exogenous chromosomal DNA fragments, such as resistance to antibiotics coded onto the vector DNA. For example, in a specific case in which pBR322 of the EK system is used as a vector, and a HindIII fragment of chromosomal DNA is inserted into the Hindlil cleavage

site of pBR322, the tetracycline resistant gene is inactivated, so a primary selection may be conducted by growth of the transformants that confer ampicillin resistance without having a HindIII cleavage site in the ampicillin gene.

Subsequently, the selected transformants are transferred onto agar plates containing starch, using, for example, a replica method, and are then cultured so as to form colonies. By staining the starch contained in the starch-containing agar plates using an iodine-containing solution, target recombinant microorganisms can be selected as they decompose starch around the colonies.

The recombinant DNA molecule harbored by the thusobtained recombinant microorganism can be extracted using standard procedures for preparing plasmids or phage DNAs (Maniatis, T. et al., Molecular Cloning, Cold Spring Harbor Laboratory, New York (1982)). When cleavage patterns obtained through the use of various restriction enzymes are analyzed by electrophoresis, it is confirmed that the recombinant DNA molecule is a ligated product of the vector DNA molecule and a DNA fragment containing the alkaline liquefying α-amylase gene.

The gene encoding an alkaline liquefying a-amylase is contained in a DNA fragment of about 2.1 kb shown in the restriction enzyme map of Fig. 1, and is present in the segment of about 1.6 kb shown by the white bar. The gene has a nucleotide sequence shown as Sequence No. 2. In this sequence, the 5' terminus and 3' terminus correspond to the left-hand side and the right-hand side, respectively, of

the fragment of about 2.1 kb shown as Sequence 2. In this sequence is observed an open reading frame (ORF) starting at the 145th nucleotide, ATG, and coding for a sequence consisting of 516 amino acid residues described in Sequence No. 1. Thirteen bases (13 b) upstream of the ORF, there exists a sequence AAGGAG which is highly complementary to the 3' terminal sequence of the 16S ribosomal RNA of Bacillus subtilis (McLaughlin, J.R. et al., J. Biol. Chem., 256, 11283 (1981)). On a further upstream region extending nucleotides from 9 to 36, there exists a sequence TTGAAA 16b TATGGT which has high homology with the consensus sequence of a oA-type promoter (Gitt, M.A. et al, J. Biol. Chem., 260, 7178 (1985)). Similarly, another oh-type promoter sequence is found at nucleotides from 95 to 125. The amino acid sequence of the 10 amino acid residues on the amino terminus side in an alkaline liquefying a-amylase purified from a culture of Bacillus sp. KSM-AP1378 coincides with the sequence extending from the 37th amino acid (amino acid Nos. 37-46 in Sequence No. 2) deduced from the nucleotide sequence of the present DNA fragment.

When the nucleotide sequence of the gene of the present invention and a deduced amino acid sequence were compared with those of a-amylase known hitherto, it was confirmed that the present gene includes a novel nucleotide sequenced, with the deduced amino acid sequence encoded by the gene being different from those of other a-amylases such as a liquefying a-amylase produced by Bacillus amylolique (Takkinen, K. et al., J. Biol. Chem., 258, 1007 (1983)), a liquefying a-amylase

produced by Bacillus stearothermophilus (Nakajima, R. et al., J. Bacteriol., 163, 401 (1985)), a liquefying α-amylase produced by Bacillus licheniformis (Yuuki, T et al., J. Biochem., 98, 1147 (1985)), or a liquefying α-amylase produced by Bacillus sp. 707 (Tsukamoto, A. et al., Biochem. Biophys. Res. Commun., 151, 25 (1988)).

An example of a preferred recombinant DNA molecule containing the entire region of the alkaline liquefying α-amylase gene is plasmid pAML100 (Fig. 2). This recombinant plasmid has a size of 4.4 kb and formed of a fragment containing a 1.8 kb fragment which contains the alkaline liquefying a-amylase gene and pUC19. An example of a preferred recombinant microorganism harboring the recombinant DNA molecule is an E. coli HB101(pAML100) strain. This strain was obtained by transforming E. coli HB101 strain with the recombinant plasmid pAML100 using a standard transformation method. When this strain is cultured using a medium routinely employed for culturing E. coli, it produces an alkaline liquefying a-amylase. The optimum reaction pH of the thus-produced enzyme is pH 8-9. This agrees well with the activity-pH relationship profile determined for the alkaline liquefying a-amylase produced by the gene donor bacterial strain, Bacillus sp. KSM-AP1378 (Fig. 4).

The DNA fragments of the present invention are not necessarily limited only to those encoding the amino acid sequences shown in the below-described sequence listing, so far as they encode a protein exhibiting the enzymatic activity of interest, and they encompass DNA fragments

encoding an amino acid sequence in which one or more amino acids are substituted, added, deleted, inverted, or inserted. An example of such DNA is one encoding an amino acid sequence equivalent to the amino acid sequence described in Sequence No. 1 from which up to 32 amino acids on the N-terminal side have been deleted.

In order to produce an alkaline liquefying α -amylase using the transformed microorganism of the present invention, a transformed microorganism harboring the aforementioned DNA fragment of the present invention is subjected to culturing. Alternatively, the DNA fragment may be integrated in a variety of expression vectors to obtain transformed microorganisms with enhanced expression ability, followed by culturing of the resultant transformants. Moreover, the transformed microorganisms may be cultured under different conditions depending on the identity of the microorganisms. Thus, culture conditions suited for the host may be used. In order to collect an alkaline liquefying α -amylase from the resultant culture, a routine method (such as the method described in W094/26881) may be used.

The DNA fragments of the present invention may be further used as probes for the isolation of homologous alkaline liquefying a-amylase genes from other organisms.

Examples

The present invention will next be described in more detail by way of examples, which should not be construed as limiting the invention thereto. Concentrations in the

Examples are all on a basis of % by weight.

Example 1:

Bacillus sp. KSM-AP1378 producing an alkaline liquefying a-amylase was inoculated in 5 ml of medium A (Table 1) and subjected to shaking culture at 30°C for 24 hours.

One ml of the culture was inoculated in 100 ml of the same medium, followed by shaking culture at 30°C for a further 12 hours. Subsequently, cells were collected by centrifugation and about 1 mg of chromosomal DNA was obtained in accordance with a method proposed by Saito and Miura (Saito, H.; and Miura K., Biochim Biophys. Acts, 72, 619 (1963)).

Table 1
Composition of medium A

Soluble starch	1.0%
Polypepton	1.0%
Yeast extract	0.5%
кн ₂ РО ₄	0.28
Na2HPO4·1ZHZO	0.25%
мgs0 ₄ ·7н ₂ 0	0.02%
CaCl _Z ·2H _Z O	0.02%
FeSO ₄ ·7H ₂ O	0.001%
MnCl ₂ ·4H ₂ O	0.0001%
Na ₂ CO ₃	1.0% (separately
	sterilized

Example 2:

It is known that many members of the amylase family

possess I-IV regions where amino acid sequences are conserved at a high level (Nakajima, R. et al., Appl. Microbiol. Biotechnol., 23, 355 (1986)). Therefore, primers 1 and 2 (Figs. 1 and 3) corresponding to regions II and IV were synthesized based on the amino acid sequence of region II and the amino acid sequence of region IV, which are particularly conserved regions among regions I through IV of known alkaline liquefying a-amylases. Using the thus-synthesized primers and chromosomal DNA of KSM-AP1378 (which served as template), PCR was conducted (one cycle = 94°C x 1 min. + 42°C x 1 min. + 60°C x 2 min., 30 cycles). A gene fragment of approximately 0.3 kb (fragment A) shown in Fig. 1 was obtained, and the nucleotide sequence of this fragment was determined. As a result, it was found that the present fragment was coded with an amino acid sequence exhibiting a non-negligible level of homology with the amino acid sequence extending from region II through region IV of known liquefying amylase.

Example 3:

Using fragment A as a probe, chromosomal DNA of Xbal-digested KSM-AP1378 was subjected to Southern hybridization. As a result, it was confirmed that there was a band which hybridized at the location of approximately 1.0 kb. An amplified fragment of approximately 0.7 kb (fragment B) was obtained by an inverse PCR method (Triglia, T. et al., Nucleic Acids Res., 16, 81 (1988)) using primers synthesized from the terminal sequences of fragment A (on the

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side of region II: primer 3; on the side of region IV: primer 4) and DNAs which had been obtained by intramolecularly ligating Kbal-digested KSM-AP1378 chromosomal DNA (Fig. 1) as template. The nucleotide sequence of fragment B was determined, which revealed that the present fragment contained a stretch, approximately 0.6 kb region downstream from region IV. The present fragment contained a termination codon for the ORF, which was deduced to be attributed to alkaline liquefying α -amylase.

Example 4:

A primer was designed and synthesized based on the N-terminal amino acid sequence (7 amino acids) of alkaline liquefying a-amylase from the KSM-AP1378 strain (Fig. 3). Using the resultant primer (primer 5) in combination with the aforementioned primer 3 (Fig. 3) and, as a template, chromosomal DNA of KSM-AP1378, PCR was conducted to obtain a fragment of approximately 0.7 kb (fragment C, Fig. 1), thereby determining its nucleotide sequence.

Example 5:

A primer containing 21 bases, stretching directly downstream of the nucleotide sequence encoding N-terminal amino acid sequence of the purified enzyme, was synthesized (primer 6). Using primers 6 and 7 (Figs. 1 and 3) and DNAs which had been obtained by intramolecularly ligating HindIII-digested KSM-AP1378 Chromosomal DNA (Fig. 1) as templates.

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fragment in which an upstream 0.8 kb fragment (fragment D) and a downstream PstI-HindIII 0.4 kb fragment had been ligated at the HindIII site. The nucleotide sequence of the fragment D region was determined, which revealed the presence of a signal sequence composed of 31 amino acids,

MKLHNRIISVLLTLLLAVAVLFPYMTEPAQA (from No. 1 to No. 31 of Sequence No. 2), a deduced SD sequence composed of AAGGAG (nucleotides 127-132; McLaughlin, J.R. et al., J. Biol.

Chem., 260, 7178 (1985)), and two kinds of deduced promoter sequences (-35 sequences, TTGAAA; -10 sequence, TATGGT, and -35 sequence, TTGACT; -10 sequence, TAAATT).

Example 6:

Using primer A located at approximately 0.1 kb upstream of the promoter sequence, primer B located 79 b downstream of the termination codon, and chromosomal DNA of KSM-AP1378 as templates, a stretch of approximately 1.8 kb between the primers was amplified by PCR. The resultant amplified fragment was inserted into the Smal site of pUC19, and then introduced into E. coli HB101. The transformant was allowed to grow on an LB agar medium containing 0.4% Starch azure and 15 µg/ml ampicillin. Colonies which had formed transparent halos around them were isolated as an E. coli strain that produced liquefying a-amylase. A recombinant plasmid was prepared from this transformant, and a restriction enzyme map of the plasmid was made. In the map, it was confirmed that an approximately 1.8 kb DNA fragment (fragment E) shown in Fig. 1 was contained. This recombinant plasmid was designated plasmid

pAML100 (Fig. 2).

Example 7:

The recombinant E. coli obtained in Example 6 was subjected to shaking culture for 12 hours in 5 ml of an LB liquid medium containing 50 µg/ml of ampicillin. One (1) ml of the culture was inoculated to 100 ml of an LB medium (containing ampicillin), followed by shaking culture at 37°C for 24 hours. Cells collected by centrifucal separation were suspended in Tris-HCl buffer (pH 8.0), and were disrupted by sonication. After the cells were sonicated, cell debris was removed by centrifugal separation, and the resultant supernatant was used as a cell-free extract. As a control, the cell-free extract of HB101(PUC19) strain was separately prepared in a similar manner. α -Amylase activities in these extracts were measured by first causing a reaction, at 50°C for 15 minutes, in a reaction mixture containing 50 mM glycine-NaCl-NaOH buffer (pH 10) and soluble starch, and then by quantitatively determining the produced reducing sugar by the 3,5-dinitrosalicylic acid method (W094/26881). One unit of enzymatic activity was defined as the amount of protein that produced a quantity per minute of reducing sugar equivalent to 1 pmol of glucose. As a result, a-amylase activity was detected in the cell-free extract of strain HB101(pAML100). The optimum working pH of α -amylase was found to fall within the pH range between 8 and 9. This result coincides well with the optimum pH of liquefying a-amylase produced by Bacillus sp. KSM-AP1378

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(Fig. 4). For the measurement of enzymatic activities, the buffers shown in Table 2 below were used (each at 40 mM).

Table 2

pH 3.5-5.5: Acetate buffer

pH 5.5-8.5: Tris-maleic acid buffer

pH 8.5-10.5: Glycine-NaCl-NaOH buffer

pH 10.5-11.0: Na₂CO₃-NaHCO₃ buffer

Industrial Applicability:

According to the present invention, it is possible to obtain a gene encoding for alkaline liquefying α -amylase exhibiting the maximum activity in the alkaline pH range as well as a microorganism harboring such gene. Use of them facilitates mass production of alkaline liquefying α -amylase.

Sequence Listing

Information for Sequence No. 1: (i) Sequence Characteristics: (A) Length: 516 amino acids (B) Type: amino acid (D) Topology: linear (ii) Molecule Type: peptide (xi) Sequence Description: Sequence No. 1: Met Lys Leo His Asn Arg Ile Ile Ser Val Leo Leo Thr Leo Leo Leo Ala Val Ala Val Leu Phe Pro Tyr Met Thr Glu Pro Ala Gln Ala His His Asn Gly Thr Asn Gly Thr Met Met Glo Tyr Phe Glu Trp His Leu Pro Asn Asp Gly Asn His Trp Asn Arg Leu Arg Asp Asp Ala Ala Asn Leu Lys Ser Lys Gly lle Thr Ala Val Trp lle Pro Pro Ala Trp Lys Gly Thr Ser Gin Asn Asp Val Gly Tyr Gly Ala Tyr Asp Leu Tyr Asp Leu Gly Glo Phe Asn Glo Lys Gly Thr Val Arg Thr Lys Tyr Gly Thr Arg Ser Gin Leu Gin Gly Ala Val Thr Ser Len Lys Asn Asn Gly He Glo Val Tyr Gly Asp Val Val Met Aso His Lys Gly Gly Ala Asp Gly -140The Glu Net Val Asn Ala Val Glu Val Asn Arg Ser Asn Arg Asn Gin

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	29	Q					295			, , , ,	, , %	300	uòn	ush	ren	Ala
Ala		g	Glu	Asn	Tyr	Leu		Lvs	Thr	Sec	Tro	Asa	u:	r>	25 🔻	n:
305						310		,	7 3	00;		13213	818	30r		
Asp	٧a	! !	oro.	Leu	His	-	Ásn	l.pu	Tur	han	315	Ser	a	n		320
					325		·······	200	121		nia	net	ASA (Gly
Tur	nı.		1							330				2	335	

Tyr Phe Asp Met Arg Asn He Leu Asn Gly Ser Val Val Gln Lys His

340

345

360

360

375

360

360

360

370

375

370

375

380

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Ala Leu lle Lou Thr Arg Glu Gln Gly Tyr Pro Ser Val Phe Tyr Gly 385 390 395 400 Asp Tyr Tyr Gly lle Pro Thr His Gly Val Pro Ser Wet Lys Ser Lys 405 410 415 He Asp Pro Leu Leu Glo Ala Arg Glo Thr Tyr Ala Tyr Gly Thr Glo 420 425 430 His Asp Tyr Phe Asp Ris His Asp Ile Ile Gly Trp Thr Arg Glu Gly 435 440 445 Asp Ser Ser His Pro Asm Ser Gly Leu Ala Thr Ile Met Ser Asp Gly 450 455 460 Pro Gly Gly Asn Lys Trp Net Tyr Val Gly Lys His Lys Ala Gly Gln 465 470 475 480 Val Trp Arg Asp lie Thr Gly Asm Arg Ser Gly Thr Val Thr lie Asm 485 490 495 Ala Asp Gly Trp Gly Asm Phe Thr Val Asm Gly Gly Ala Val Ser Val 500 505 510 Trp Val Lys Glo 516

Information for Sequence No. 2:

- (i) Sequence Characteristics:
 - (A) Length: 1775 base pairs
 - (B) Type: nucleic acid
 - (C) Strandedness: double
 - (D) Topology: linear
- (ii) Molecule Type: DNA (genomic)
- (vi) Original Source:
 - (A) Organism: Bacillus sp.

(B) Strain: KSM-AP1378

(xi) Sequence Description: Sequence No. 2:

								iA TA									60
								TT									120
AAA	TTGA	AGG	AGAG	GGTG	CT T	TTT	ATG	AAA .	CTT	CAT	AAC	CGT	ATA	ÀŦŤ	AGC	GTA	174
							Met	Lys	Leu	His	Åsn	Arg	He	He	Ser	Val	
							1				S					10	
CTA	TTA	ACA	CTA	TTG	TTA	GCT	GTA	GCT	611	TTG	TTT	CCA	TAT	AT6	ACE	° \$	222
Leu	Leu	Thr	Leu	Leu	Leu	Ala	¥a]	Ala	Val	Leu	Pho	Pro	Tyr	Wet	Thr	•	
				15					20					25			
GAA	CCA	GCA	CAA	GCC	CAT	CAT	AAT	666	ACG	AAT	666	ACC	ATG	ATG	CAG	`	270
								Gly									
			30					35					40				
TAT	TŤT	GAA	T66	CAT	776	CCA	AAT	GAC	GGG	AAC	CAC	TGG	AAC	AGG	TTA		318
								Àsp									
		45					50					55					
CGA	GAT	GAC	GCA	GCT	AAC	TTA	AAG	AGT	AAA	666	ATT	ACC	GCT	GTT	766		366
								Ser									
	60					65					70						
ATT	CCT	CCT	GCA	TGG	AAG	666	ACT	TCG	CAA	AAT	GAT	GTT	666	TAT	661		414
lle	Pro	Pro	Ala	Trp	Lys	Gly	Thr	Ser	61n	Aso	Asp	Val	Gly	Tyr	Gly		
75					80					85					90		
GCC	TAT	GAT	TTG	TAC	GAT	CTT	GGT	GAG	TTT	AAC	CAA	AAG	GGA	33A	GTC		462
								Glu									
				95					100					105			
CGT	ACA	AAA	TAT	GGC	ACA	AGG	AGT	CAG	TTG	CAA	661	GCC	676	ACA	TU		510

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			110					115					120			
TTC	i AAA	raa i	" AAC	GGC	ATI	°, CAA	CTI	TAT	` 6 66	GAT	GT(org	ATO	AAT	CAT	558
tai	i lys	Asr	Asn	Gly	116	61n	¥a1	Tyr	Gly	Asp	Val	. Val	Met	Asn	llis	
		125					130					135				
															AAC	808
Lys	Gi y	Gly	Ala	Åsp	Gly	Thr	Glu	Met	Val	Åsn	Ala	Val	Glu	Val	Às'n	
	140					145					150		·			
CGA	AGC	AAE	CGA	AAC	CAA	GAA	ATA	TCA	eet	GAA	TAC	ACC	ATT	GAA	SCA	654
Årg	Ser	Asn	Arg	Asn	Gln	Glu	lle	Ser	Gly	Glu	Tyr	Thr	lle	Glu	Ala	
155					160					165					170	
TGG	ACG	AAA	TTT	GAT	TTC	CCT	GGA	AGA	GGA	AAT	ACC	CAT	TCC	AAC	TTT	702
			Phe													
				175					180					185	•	
ÅÅÅ	TGG	CGC	TGG	TAT	CAT	TTT	GAT	GGG	ACA	GAT	T66	GAT	CAG	TCA	CGT	750
			Trp													
			190					195					200			
CAG	CTT	CAG	AAC	AAA	ATA	TAT	AAA	TTC	AGA	GGT	ACC	66A	AAG	SCA	TGG	798
			Åsn													
		205					210		41			215			r	
GAC	TGG	GAA	GTA	GAT	ATA	GAG.	AAC	66C	AÁC	TAT	GAT	TAC	CIT	ATG	TAT	846
			Val													~
	220					225					230				·	
GCA	GAC	ATT	GAT	ATG	GAT	CAT	CCA	GAA	STA	ATC	AAT	GAA	CTT	AGA	AAT	894
			Asp													
235					240					245					250	
TGG	GGA	GTT	TGG	TAT	ACA	AAT	ACA	CTT	AAT	CTA	GAT	GGA	TTT	AGA		942
			Trp													**************************************
				255					260		·	*		285	V . M	
														and the Ph		

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GAT	SET	GTG	AAA	CAT	ATT	ÀÀÁ	TAC	AGC	TAT	ACG	AGA	GAT	760	CTA	άCλ	990
Asp	Ala	Val	(ys	1115	lle	Lys	lat	Sec	Tyr	Thr	Arg	yab	Trp	Leu	Thr	
			270					275					280			
TA3	GTG	CGT	AAC	ACC	ACA	667	AAA	CCA	ATG	777	GCA	GTT	GCA	GAA	TTT	1038
His	Val	Arg	Asn	Thr	Thr	Gly	Lys	Pro	Net	Phe	Ala	Val	Ala	Gtu	Phe	
		285					290					295				
7,66	AAA	AAT	SAC	CTT	GCT	GEA	ATC	GAA	AAC	TAT	TTA	TAA	AAA	ACA	AGT	1088
Trp	l.ys	Àsn	Asp	Leu	Ala	Ala	118	Stu	Åso	Tyr	Leu	Àsn	Lys	Thr	Ser	
	300					305					310					
TGG	AAT	CAC	TEC	GTG	TTC	GAT	GTT	133	TIT	CAT	TAT	AAT	TTG	TAC	AAT	1134
Trp	Asn	His	Ser	Val	Phe	Asp	Val	pro	Leu	His	Tyr	Asn	Leu	Tyr	Aşn "	
315					350					325					330	
GCA	TET	AAT	AGT	GGT	GGC	TAT	TTT	GAT	ATG	AGA	AAT	ATT	TTA	AAT	GGT	1182
Ala	Ser	Asn	Ser	Gly	Gly	Tyr	Phe	Åsp	Met	Arg	Asn	He	Leu	Asn	Gly	
				335					340					345		
			CAA													1230
Ser	Va 1	Val	Gln	Гàв	His	Pro	He	His	Ala	Val	Thr	Phe	Val	Åsp	Asn	
			350					355					360			
CAT	GAC	TCT	CAS	CCA	GGA	GAA	GCA	776	GAA	TCC	777	GTT	CAA	TCG	TGG	1278
His	Asp	Ser	610	510	Gly	Glu	Ala	Leu	Glu	Ser	Phe	Val	Gln	Ser	Trp	
		365					370					375				
TTC	AAA	CCA	CTG	GCA	TAT	GCA	TTG	ATT	CTG	ACA	AGG	GAG	CAA	GGT	TAC	1326
Phe	Lys	Pro	Leu	Ala	Tyr	Ala	Leu	He	Leu	Thr	Arg	Glu	Gln	Gly	Tyr	
	380					385					390					
CCT	TCC	GTA	TTT	TAC	661	GAT	TAC	TAC	GGT	ATA	CCA	ACT	CAT	GGT	GTT	1374
Pro	Ser	Val	Phe	Tyr	Gly	Asp	Tyr	Tyr	Gly	He	Pro	The	llis	Gly	Val	
- 395					400					405					410	
CCT	TCG	ATG	۸۸۸	TCT	۸۸۸	ATT	GAT	CCA	CIT	CTG	CAG	GCA	CGT	CAA	ACG	1422
Pro	Ser	Met	Lys	Sec	Lys	He	Åsp	Pro	Leu	Lés	Gin	Ala	Arg	Gla	Thr	

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				415					420					425			
TAT	GCC	TAC	GGA	ACC	CAA	EAT	GAT	TAT	TIT	GAT	CAT	CAT	GAT	ATT	ATC	1.4	170
Tyr	Ala	Tyr	Gly	Thr	Gla	His	Asp	Tyr	Phe	Asp	llis	llis	Ásp	He	lle		
			430					435					440				
660	TGG	ACG	AGA	GAA	666	GAC	AGC	TCC	CAC	CCA	AAT	TCA	GGA	CTT	GCA	18	18
Gly	Trp	Thr	Arg	Glu	Gly	Asp	Ser	Ser	His	Pro	Asa	Ser	Gly	Lea	Ala		
		445					450					458	š				
ACT	ATT	ATG	TCC	GAT	666	CCA	666	SGT	AAT	AAA	TGG	ATG	TAT	STC	GGG	15	66
The	He	Met	Ser	Asp	Gly	Pro	Gly	Gly	Asn	Lys	Trp	Net	Туг	Val	61y		
	460					465					470						
AAA	CAT	AAA	GET	GGC	CAA	GTA	TGG	AGA	GAT	ATC	ACC	GGA	AAT	AG6	TCT	16	14
Lys	His	Lys.	Ala	Gly	61n	Val	Trp	Arg	Asp	lle	Thr	Gly	Asn	Arg	Ser		
475					480					485					490		
GGT	ACC	GTC	ACC	ATT	AAT	GCA	GAT	SGT	TGG	666	AAT	TTC	ACT	GTA	AAC	16	62
Gly	Thr	Val	Thr	Ne	Àsņ	Ala	Asp	Gly	Tro	Gly	Àsn	Phe	Thr	Va1	Asn		
				495					500					505			
GGA	GGG	GCA	GTT	TCG	GTT	TGG	GTG	AAG	CAA	TAAA	TAA6	iga /	CAAC	SAGGE	36	17	12
Gly	Gly	Ala	Val	Ser	Val	Trp	Val	Lys	Gin								
			510					515									
AAA	NTTA(TT 1	ract/	CATO	ac as	SAGCT	TTC	. GAT	CACT	CAT	ACAC	CCA/	YTA 1	ľaaa"	TTGGA	A 17	72
GCT																	76

CLAIMS:

1. A DNA molecule encoding alkaline liquefying α -amylase activity.

- Z. A DNA molecule as defined in Claim 1, which encodes the amino acid sequence described in Sequence No. 1 or a functional fragment thereof.
- 3. A DNA molecule encoding a protein exhibiting alkaline liquefying a-amylase activity and possessing an amino acid sequence described in Sequence No. 1 in which one or more amino acids are substituted, added, deleted, inverted, or inserted.
- 4. A DNA molefule as defined in any one of Claims 1 through 3, further comprising a nucleotide sequence for regulating expression of a gene.
- 5. A recombinant DNA containing the DNA molecule of any one of Claims 1 through 4.
- 6. A transformed microorganism harboring the recombinant DNA of Claim 5.
- 7. A method for producing alkaline liquefying a-amylase, comprising culturing the transformed microcrganism of Claim 6 and isolating the alkaline liquefying a-amylase produced by the microorganism.
- 8. A DNA molecule which hybridizes to a DNA sequence which is complementary to the nucleic acid sequence of SEQ ID No. 2.
 - 9. A protein encoded by the DNA molecule of Claim 9.
- 10. A DNA molecule which hybridizes to a DNA sequence which is complementary to the nucleic acid sequence of SEQ ID No. 2, wherein said DNA molecule encodes a protein having alkaline

liquefying a-amylase activity.

11. A protein encoded by the DNA molecule of Claim 11.

- 12. The recombinant DNA plasmid pAML100.
- 13. The recombinant E. coli strain HB101(pAML100).

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The present invention provides a DNA fragment encoding alkaline liquefying a-amylase, recombinant DNA containing the DNA fragment, a transformed microorganism harboring the recombinant DNA, as well as a method for producing alkaline liquefying a-amylase using the transformant. The method of the present invention enables mass production of alkaline liquefying a-amylase useful as a detergent component.

AMENDED CLAIMS

[received by the International Bureau on 11 December 1996 (11.12.96); original claims 4, 9, 11 amended; remaining claims unchanged (2 pages)]

- 1. A DNA molecule encoding alkaline liquefying α -amylase activity.
- 2. A DNA molecule as defined in Claim 1, which encodes the amino acid sequence described in Sequence No. 1 or a functional fragment thereof.
- 3. A DNA molecule encoding a protein exhibiting alkaline liquefying a-amylase activity and possessing an amino acid sequence described in Sequence No. 1 in which one or more amino acids are substituted, added, deleted, inverted, or inserted.
- 4. A DNA molecule as defined in any one of Claims 1 through 3, further comprising a nucleotide sequence for regulating expression of a gene.
- 5. A recombinant DNA containing the DNA molecule of any one of Claims 1 through 4.
- 6. A transformed microorganism harboring the recombinant DNA of Claim 5.
- 7. A method for producing alkaline liquefying α -amylase, comprising culturing the transformed microorganism of Claim 6 and isolating the alkaline liquefying α -amylase produced by the microorganism.
- 8. A DNA molecule which hybridizes to a DNA sequence which is complementary to the nucleic acid sequence of SEQ ID No. 2.
 - 9. A protein encoded by the DNA molecule of Claims 1 through 4.
- 10. A DNA molecule which hybridizes to a DNA sequence which is complementary to the nucleic acid sequence of SEQ ID No. 2, wherein said DNA molecule encodes a protein having alkaline

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liquefying a-amylase activity.

11. A protein encoded by the DNA molecule of Claim 10.

- 12. The recombinant DNA plasmid pAML100.
- 13. The recombinant E. coli strain H8101(pAML100).

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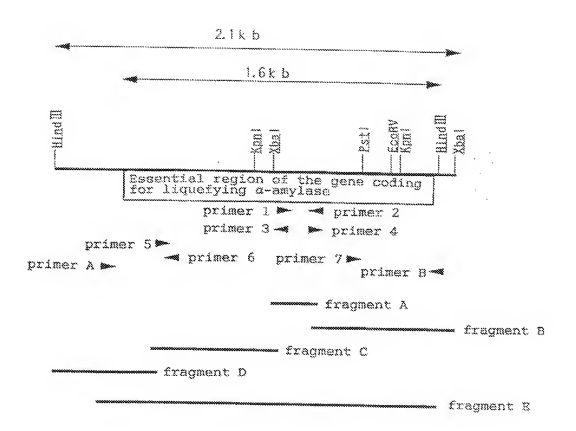


FIG. 1

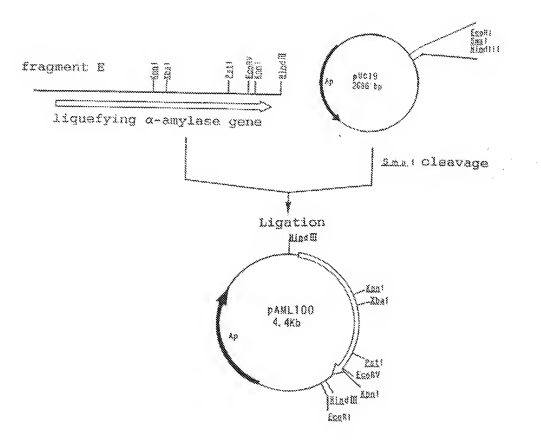


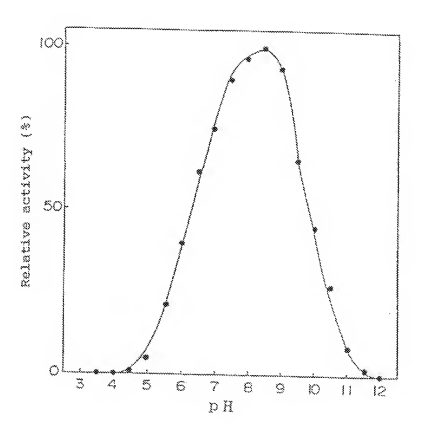
FIG. 2

FIG. 3

primer 1 5 TAGACGCAGTAAAACACATAAA 3 CTCCGTC 6 G G TT Ţ 3' CGACAATGAAAACAACTATTAGTACT 5' primer 2 primer 3 5° AGCCAATCTCTCGTATAGCTGTA 3° primer 4 5° GTACAAAAACACCCTATACATG 3° primer 5 5° AATGGAACAATGATGCAGTA 3° primer 6 5' CATTIGGCAAATGCCATTCAAA 3' primer 7 5'AAAATTGATCCACTTCTGCAG 3' primer A 5° CAGCGCGTGATAATATAAATTTGAAT 3° 5° AAGCTTCCAATTTATATTGGGTGTAT 3° primer B

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FIG. 4



INTERNATIONAL SEARCH REPORT

tests: used Appropriation No PCT/JP 96/01641

PCT/JP 96/01641 A. CLASSIFICATION OF SOMECT MATTER IPC 6 C12N15/56 C12N9/28 C12N1/21 C12N15/70 According to International Patent Conflication (IPC) or to both national classification and (PC B. PIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) 1PC 6 CIZN Documentation removed other than minimum documentation to the extent that such documents are included in the fields respected Electronic data base consisted during the international search (name of data have and, where practical, search terms used) C. DOCUMENTS CÓNSIDERED TO BE RELEVANT Category ' Citation of document, with indication, where appropriate, of the retevant passages Externat to claus, too, X WO.A.94 26881 (KAO CORP :ARA KATSUTOSHI 1-13 (JP); SAEKI KATSUHISA (JP): IGARASHI KAZU) 24 November 1994 cited in the application see the whole document & EP.A.O 670 367 (KAO CORPORATION) Х BIOCHEMICAL AND BIOPHYSICAL RESEARCH 1,3-11 COMMUNICATIONS. vol. 151, no. 1, 29 February 1988, pages 25-31, XP000605386 TSUKAMOTO A. ET AL.: "Nucleotide sequence of the maltohexaose-producing amylase gene from alkalophilic Bacillus sp. #707 and structural similarity to liquefying type alpha-amylases." A see the whole document 2,12,13 -/--X Further documents are listed in the communication of hox C. X Patent family merehers are fisted in somex. * Special categories of cated documents: "I" later document published after the marmational filing date or priority date and not in combine with the application bin vised to trademand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of puritoidar relevance. "E" earlier document but published on or after the international "X" document of particular retevance; the claimed inventors cannot be considered povel or cannot be considered to Ming date "E" document which may throw doubts on priority claims) or mustbe an inventive step when the document is taken alone which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to myoles an inventive step when the focument is combined with one or more other such docudocument referring to an oral disclosure, use, exhibition or other means ments, such combination being obvious to a person skilled document published prior to the international filling date but later than the priority date claimed "&" deciment member of the tame patent family Date of the actual completion of the international search Date of mailing of the international search report 15. 11. 96 6 November 1996 Name and mailing address of the ISA Authorized officer Europeas Patent Office, P.B. 3818 Patentiaen 2 N.L. 2330 HV Birmsig Tel. (* 31-75) 340-2540, Tx. 31 651 epo st., Pan (* 31-75) 340-2540, Mandl. B

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INTERNATIONAL SEARCH REPORT

Inter and Application So PCT/JP 96/01641

***************************************		PCT/JP 96/01641
Confinu Tegnty	asson) DOCUMENTS CONSIDERED TO BE RELEVANT Classon of document, with indication, where appropriate, of the relevant passages	Eslevent to claum 350.
(JOURNAL OF BIOCHEMISTRY, vol. 98, no. 5, 1985, pages 1147-1156, XP002017641 YUUKI T ET AL.: "Complete nucleotide sequence of a gene coding for heat- and pH-stable alpha-amylase of Bacillus licheniformis: Comparison of the amino acid sequences of three bacterial liquefying alpha-amylases deduced from the	1,3-11
\	DNA sequences." see the whole document	2,12,13
4	EP,A.0 410 498 (GIST BROCADES NV ;PLANT GENETIC SYSTEMS NV (BE)) 30 January 1991 see the whole document	3
Ρ,Χ	w0,A,95 26397 (NOVONORDISK AS ;OUTTRUP HELLE (DK); BISGAARD FRANTZEN HENRIK (DK);) 5 October 1995 see the whole document	1,3-11
	-	

INTERNATIONAL SEARCH REPORT

information on patent family members

Inter onal Application No PCT/JP 96/01641

Patent document cited in search report	Publication date	Patent memi	Publication date		
WO-A-9426881	24-11-94	CN-A- EP-A-	1110058 0670367	11-10-95 06-09-95	
EP-A-0410498	38-01-91	AU-B- AU-A- CA-A- CN-A- WO-A- JP-T- US-A-	638263 5953890 2030554 1050220 9100353 4500756 5364782	24-06-93 17-01-91 30-12-90 27-03-91 10-01-91 13-02-92 15-11-94	
W0-A-9526397	05-10-95	AU-A- ZA-A-	2067795 9502565	17-10-95 21-12-95	